RECEIVED #3

FORM PTO-1449

ORMATION DISCLOSURE CITATION

Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to Applicant.

JUN 0 6 7800 1 of 21 Attorney Docket Serial Number 22789XA-TTECH CENTER 4660 29007

Applicant

STEINER et al.

Filing Date Feb. 13, 2001

Group Art Unit 1614

	Feb. 13, 2001 1614									
U.S. PATENT DOCUMENTS										
Examiner Initial		Document Number	Date		Name	Class	Sub- Class	Filing Date		
\mathcal{N}	AA	5,532,248	7/2/96	Gou	let et al.	1		5/12/95		
	AB	5,506,228	4/9/96	Nort	on et al.			2/23/95		
	AC	5,470,878	11/28/95	Michr	nick et al.			12/8/93		
	AD	5,457,111	10/10/95	Lu	ly et al.		/	11/9/93		
	AE	5,385,918	1/31/95	Conn	ell et al.		X	2/9/93		
	AF	5,342,625	8/30/94	Hau	er et al.			12/15/92		
	AG	5,292,747	3/8/94	Dav	is et al.			9/21/92		
	AH	5,284,877	2/8/94	Org	an et al.			6/12/92		
	ΑI	5,284,840	2/8/94	Ruppr	echt et al.			6/12/92		
	AJ	5,284,826	2/8/84		Eberle			8/27/92		
AK 5,258,389		11/2/93	Goul	et et al.	1/		11/9/92			
FOREIGN PATENT DOCUMENTS										
		Document Number	Date	(Country	Class	Sub- Class	Trans- lation		
V	ΑL	₩ 0, 9641609	12/27/96	S	PCT			Yes		
	AM	FP 564924	10/13/93	ÉΡ	#P			Yes		
	AN	₽ 423714	4/24/91	G P	Y K			Yes		
n'/	AO	∜ O 9736869	10/9/97	W	pdr			Yes		
						<u> </u>				
	OTHER	(Including Au	thor, Tit	le, Dat	e, Pertinent	Pages,	etc.)			
n	AP	Birkensha and Synth Medicinal	w, T.N. e esis of P Chemistr	t al., yranose <u>y Lette</u>	"Synthetic FK Replacements <u>rs</u> , (1994) 4:	BP12 L s," <u>Bio</u> 21, 25	igands <u>organi</u> 01-250	. Design <u>c &</u> 6.		
	AQ	Caffrev.	M.V. et a	1 "Sv	nthesis and E Ligands," <u>Bi</u> 4:21, 2507-2	valuat	ion of	Dual		
n	AR	Hauske, J	R et al	. "Desi	gn and Synthe nal Chemistry	sis of	Novel	FKBP		
Examiner	/	2 Certs			Date Consider	ed				

Attorney Docket Serial Number 22789XA-T 09/781,427 FORM PTO-1449 JUN 0 4 2001 Applicant STEINER et al. MATION DISCLOSURE CITATION Filing Date Group Art Unit Feb. 13, 2001 1614 U.S. PATENT DOCUMENTS Examiner Document Sub-Filing Date Class Name Initial Number Class Date W BA 5,208,241 5/4/93 Ok et al. 9/9/91 BB 5,192,773 3/9/93 Armistead et al. 7/2/90 BC 5,189,042 2/23/93 Goulet et al. 8/22/91 BD 4,996,193 2/26/91 Hewitt et al. 3/3/89 BE 3/25/97 5,614,547 Hamilton et al. 6/7/95 FOREIGN PATENT DOCUMENTS Document. Sub-Trans-Date Country Class Number Class lation Yes No Yes No Yes No Yes No Yes No OTHER (Including Author, Title, Date, Pertinent Pages, Holt, D.A. et al., "Design, Synthesis, and Kinetic Evaluation of High-Affinity FKBP Ligands and the X-ray Crystal Structures of Their Complexes with FKBP12," <u>J. Am.</u> Chem. Soc., (1993) 115, 9925-9938. W BF Holt, D.A. et al., "Structure-Activity Studies of Synthetic FKBP Ligands as Peptidyl-prolyl Isomers Inhibitors, "Bioorganic & Medicinal Chemistry Letters, (1994) 4:2, 315-320. BG Holt, D.A. et al., "Structure-Activity Studies of Nonmacrocyclic Rapamycin Derivatives," Bioorganic & N. BHMedicinal Chemistry Letter, (1993) 3:10, 1977-1980. Cook Date Considered Examiner

EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP \$ 609.

Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to Applicant.

RECEIVED

OIP	<u> </u>			•		TE(CH CENT	ER 1000	2903 of 21	
JUN 0 4 2	yC131	FORM PTO-144	a			ey Dock 89XA-T		rial N		
JUN 0 4 2	, E. S.	10Id1 110-144	<i>3</i>		Applica		TMED -			
RADELAN	PORMA	TION DISCLOSUR	E CITATIO	<u>N</u>	STEINER et al.					
					Filing Date Group Art Uni Feb. 13, 2001 1614					
			U.S. PAT	ENT DO						
Examiner		Document	<u> </u>	<u> </u>		1		Sub-	Filing	
Initial		Number	Date		Name		Class	Class	Date	
				-						
								·		
									<u></u>	
		V. =			· · · · ·					
		I	FOREIGN PA	TENT D	OCUMENT	S				
		Document Number	Date		Country		Class	Sub- Class	Trans- lation	
									Yes No	
									Yes No	
									Yes No	
									Yes No	
									Yes No	
	OTHER	R (Including Au	thor, Tit	le, Da	te, Per	tinent	Pages	, etc.)		
N	CA	Luengo, J Relations <u>Medicinal</u>	.I. et al hips of M Chemistr	., "Syr acrocyc	thesis	and St: 3P Ligar	ructur nds,"	e-Acti Bioorg	vity anic &	
1										
	СВ	Snyder, S System,"	Nature Me	<u>dicine</u> ,	(1995)	1:1,	32-37.			
D	Teague, S.J. et al., "Synthesis and Study of a Non-Macrocyclic FK506 Derivative," <u>Bioorganic & Medicinal</u> Chemistry Letters, (1994) 4:13, 1581-1584.									
	CD	Steiner, 1997.								
Examiner	R	Cook	· -		Date Co	nsidere	ed (0/			
Draw 1	ine through	rence considered, whether or gh citation if not in confor this form with next communic	mance and not consi	idered.	with MPEP § 60		· ·			

RECEIVED

OIPE				<u> </u>		TECH	CENTER 9	6002098 21		
JUN 0 4 201	C181	HODY DEG 144	•		Attorney Dock 22789XA-T	- गम्पा	rial N			
		FORM PTO-144	9		Applicant					
RADEMAN	FORMA	TION DISCLOSUR	E CITATIO	<u>N</u>			et al.			
		FORM PTO-144			Filing Date Feb. 13, 200		roup Art	t Unit 514		
	1		U.S. PAT	ENT DO	CUMENTS					
Examine r Initial		Document Number	Date		Name	Class	Sub- Class	Filing Date		
11110101						 	_			
										
				-						
				:						
				 .	·	<u> </u>				
		I	FOREIGN PA	TENT D	OCUMENTS	I	1			
		Document Number	Date		Country	Class	Sub- Class	Trans- lation		
		in an angel						Yes No		
								Yes No		
				-		-		Yes No		
								Yes No		
							<u> </u>	Yes No		
	OTHE	R (Including Au								
n	DA	Domain of <u>& Medicin</u>	.J. et al FK-506 fo al Chemis	., "The or the <u>try Let</u>	Affinity of Immunophilin ters, (1993)	the Ex FKBP12 3:10,	cised : 2," <u>Bio</u> 1947-1	Binding organic 950.		
	Wang, G.T. et al., "Synthesis and FKBP Binding of Small Molecule Mimics of the Tricarbonyl Region of FK506," Bioorganic and Medicinal Chemistry Letters, (1994) 4:9, 1161-1166.									
N	DC	Yamashita of Dual D Chemistry	, D.S. et omain FKB , (1994)	al., " P Ligar 4:2, 32	Design, Synth ds," <u>Bioorgan</u> 5-328.	esis a	and Eva Medicin	luation al		
Examiner		R Cooli			Date Considere	ed				
Draw 1	ine throu	erence considered, whether or agh citation if not in conform f this form with next communic	mance and not cons	ldered.	• • • • • • • • • • • • • • • • • • • •					

		FORM PTO-144		Attorney Docket Serial Number 22789XA-T 09/781,427					
	TODY				Applicant S	TEINER		t al.	
<u> 1N</u>	FORMA	ATION DISCLOSUR	E CITATIO	<u>N</u>	Filing Date Feb. 13, 2			oup Ar	t Unit
			U.S. PAT	ENT DO				-	
Examiner Initial		Document Number	Date		Name	Cla	ss	Sub- Class	Filing Date
OIPE									
<u> </u>	(3)								
JUN 0 4 20	<u>\$</u>								
TRADEMAN	S						1		

		-					_		
	<u> </u>								
	I		FOREIGN PA	TENT DO	OCUMENTS				
<u> </u>		Document Number	Date	(Country	Clas	ss	Sub- Class	Trans- lation
							4		Yes No
							_		Yes No
									Yes No
	-			· · · · · · · · · · · · · · · · · · ·			\dashv		Yes No
	OMITE	D (Implication Du	m						Yes No
	OTHE	T							<u> </u>
N	EA	peptidyl- cyclospor growth in	prolyl cia in A, FK50 itiation a for hair o	s-trans 06, asc in mous	ects of immursomerase omycin and re: immunosumursumursumursumursumursumursumursumu	(PPIaso rapamyo opress	e i cin ion	inhibit n, on l n is no	nair ot
	EB	l Itopical a	pplication	n of FK	mulation of 506, a poter <u>ol</u> , (1994) 1	at immi	นิทด	suppre	essive
h	EC	l skin by to	opical apr	olicati	ion of anage on of FK506, vest. Dermat	. a not	ten	nt T	
Examiner		1 Cools			Date Conside				
Draw 1	ine throu	erence considered, whether or agh citation if not in conform this form with next communic	ance and not consi	dered.					·
Incida	- copy or	LOIM WICH HEAL COMMUNIC	acton co wbbileaut	•					

FORM PTO-1449

INFORMATION DISCLOSURE CITATION

Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to Applicant.

22789XA-T

Attorney Docket Serial Number 09/781,427

Applicant

STEINER et al.

Filing Date

					ing Date eb. 13, 200		oup Art 16	Unit			
U.S. PATENT DOCUMENTS											
Examiner InOtia		Document Number	Date	Na	ame	Class	Sub- Class	Filing Date			
ÙΥ	گاً⁄∀	5,447,915	9/5/95	Schreibe	1	/	8/28/92				
JUN 0 4 200	FΒ	5,294,603	3/15/94	Rinehar	rt, K.L.		K	2/18/92			
E.	8 7℃	5,359,138	10/25/94	Takeuch	i et al.			6/29/92			
TADEMARY.	FD	5,516,797	5/14/96	Armistea	d et al.	/		4/11/94			
FOREIGN PATENT DOCUMENTS											
		Document Number	Date	Cou	ntry	Class	Sub- Class	Trans- lation			
N	FE	₩Ф9203472	3/5/92		CT			Yes			
	FF	JE04149166	5/22/92		pan		/	No			
	FG	EP-468339	1/29/92		Þφ	<u> </u>		Yes			
	FH	W09113088	9/5/91		A T	/		Yes			
	FI	EP-405994	1/2/91		PD		1	Yes			
W	FJ	E∳-378318	7/18/90	GP E	k p			Yes			
	OTHE	R (Including Au	thor, Tit	le, Date,	Pertinent	Pages,	etc.)				
W	FK	Askin, D. versatile 55(20), 5	synthetic	"Effecient c intermed	Degradation	on of 1 Org. Cl	FK-506 hem.,	to a 1990,			
	FL	the trica	rbonyl co	ntaining m	Joshua, "De acrolide ra 34), 4845-	apamyc:	tive s in,"	tudies on			
	FM	lapplicati	on of Eva osuppress	ns technol	of tricarlogy to the	total	svnth	esis of			
	FN	Exploration Exploration	on of some	"A formal e alternat 55(9), 278	synthesis of ives to made 6-97.	of FK-! crolact	506. tamiza	tion," J.			
Rao, A.V., et al., "Studies directed towards the synthesis of immunosuppressive agent FK-506: construction of the tricarbonyl moiety," Tetrahedron Lett., 1990, 31(10), 1439-42.											
N	FP	Harding, immunosup isomerase	M.W., et pressant ," Nature	al., "A re FK506 is a Lett., 19	ceptor for cis-trans 89, 341, 7	the peption	dyl-pr	olyl			
Examiner	14	^ /	Date Considered								
	R	Civh			9/11/0	/					
EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP \$ 609.											

FORM PTO-1449 INFORMATION DISCLOSURE CITATION

Attorney Docket 22789XA-T Serial Number 09/781,427 Applicant STEINER et al. Group Art Unit 1614 Filing Date 2001 Feb. 13,

U.S. PATENT DOCUMENTS										
Examiner		Document Number	Date	Name	Class	Sub- Class	Filing Date			
N.	CA CB	4,818,749	4/4/89	Gold, E.H. et al.			4/4/89			
JUN 0 4 2001		4,808,573	2/28/89	Gold , E.H. et al.			2/28/89			
	E GC	5,252,579	10/12/93	Skotnicki, J. et al.			2/16/93			
PADEMARK	GD	5,543,423	8/6/96	Zelle et al.	7	\	1/23/95			
FOREIGN PATENT DOCUMENTS										
		Document Number	Date	Country	Class	Sub- Class	Trans- lation			
\sim	GE	₩09104985	4/18/91	MO BCI			Yes			
	GF	DE4015255	11/14/91	D6 Germany			No			
	GG	EP-419049	3/27/91	EP EPP		<u> </u>	Yes			
	GH	W09012805	11/1/90	WO PCT			No			
	GI	巨中-352000	1/24/90	EP FRO			Yes			
	GJ	DE3931051	3/29/90	by Germany			No			
\mathcal{N}	GK	EP-333174	9/20/89	Eb #40	<u> </u>		Yes			
	OTHE			le, Date, Pertinent						
~	GL	and Prese	rvation o	et al., "Prevention f CD4 Function by th 990, 249, 287-91.	of HIV e Bind:	V-1 In: ing of	fection CPFs to			
W	GM	Goodfellov and diazo Cross-Lin	w, Val S. pyruvamid king Biop	et al., "p-Nitrophe es, a New Family of robes," Biochemistry	nyl 3-0 Photoa , 28(1	diazopy ctivata 5), 634	yruvate able 46-60.			
	GN	Wasserman region of document of 54(22), 54	, H.H. et FK-506 t cited in 406.	al., "Synthesis of hrough and amidophos CA111(7):57366p}," J	the tiphorane	ricarbo e [Erro Chem.	onyl atum to , 1989,			
y	GO	Wasserman region of 1989, 54()	, H.H. et FK-506 t 12), 2785	al., "Synthesis of hrough an amidospher -6.	the tra	icarbo Org. (nyl Chem.,			
	Askin, D. et al., "Chemistry of FK-506: benzilic acid rearrangement of the tricarbonyl system," Tetrahedron Lett., 1989, 30(6), 671-4.									
h	GQ	manipulat:	ions of this	anishefsky, S., "Deg he immunosuppressant etic intermediates,"	FK506	: prepa	aration			
Examiner	Examiner $\rho_{l(lo)}$ Date Considered:									

EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPBP § 609.

Draw line through citation if not in conformance and not considered.

								<u> </u>	e 8 of 21
					:	Attorney Dock 22789XA-T	et Se	rial N 09/78	umber 31,427
INI	FORMA		M PTO-1449 DISCLOSURI		<u>N</u>	Applicant STE	INER e	t al.	
						Filing Date Feb. 13, 200	Gre	oup Ar	t Unit
				U.S. PAT	ENT DO	CUMENTS		-	
Examiner I <u>nit</u> ial			ocument Jumber	Date		Name	Class	Sub- Class	Filing Date
OME	HA	4,	574,079	3/4/86	Gavras	, H.P. et al.	,		3/4/86
JUN 0 4 2001	нв	5,	330,993	7/19/94	Armis	tead, et al.			7/2/91
	3 /	_	F	OREIGN PA	TENT D	OCUMENTS			
PADEMARK OF			ocument Jumber	Date		Country	Class	Sub- Class	Trans- lation
U	HС	MO	8809789	12/15/88	Νo	ÞУТ			Yes
	HD	ĦР	-260118	3/16/88	ÉP	H ₀			Yes
W_	HE	₩0.	9617816	6/13/96	wJ	k <i>o</i> t			Yes
	OTHER	R (In	cluding Au	thor, Tit	le, Da	te, Pertinent	Pages,	etc.)	
			The second secon		ador i dilitare de servega di jumpa per	alleggi ann agus an ann ann agus aire ann agus aire ann an agus ann ann ann an ann ann ann ann ann ann		nganamin gallandi dipelikikani	nay
, n	НF		Electroch ketoamide	emical Re s. II. El rom S-(-)	duction ectrore -prolin	'Stereochemist n of Optically eduction of be ne," Bull. Soc	Activ nzovlf	e α- ormami . Fr.,	des 1989,
~	HG		Soai, Ken amides De 1986, 24,	rived fro	m (S)-p	nmetric Allyla proline esters	tion o ," Pep	f α-ke t. Che	to m.,
	нн								
a	нЈ		Egbertson the trica: 54(1), 11	, M. and rbonyl re -12.	Danishe gion of	efsky, S., "A FK-506," J.	synthe Org. C	tic ro	ute to 1989,
	нк		Williams, diketo am FK506," J	D.R. and ide segme . Org. Ch	Benbownt of tem., 19	v, J.W., "Synt the novel immu 988, 53(191),	hesis nosupp 4643-4	of the ressiv ·	α,β e
	HL		Kocienski segment o 1988, 29(f tsukuba	enolide	synthesis of e (FK506)," Te	the C(trahed	1)-C(1 ron Le	5) tt.,
N	НМ		Tanaka, H imunosupp: Soc., 198	ressant i	solated	cture of FK506 d from Strepto L-3.	, a no myces,	vel " J. A	m. Chem.
Examiner		A	Cwh			Date Consider 9	ed ////o/	,	
Draw 1									

	FORM PTO-1449					Attorney Docket Serial N 22789XA-T 09/7			umber 81,427	
<u>IN</u>	FORMA	TION DISCLOSUR	E CITATIO	<u>N</u>	Applicant STEINER et al.					
					Filing I Feb. 1	Date 13, 200		oup Art	Unit	
			U.S. PAT	ENT DO	CUMENTS					
Examiner Initial		Document Number	Date		Name		Class	Sub- Class	Filing Date	
4	IA	4,374,829	2/22/83	Harris	, E.E. €	et al.			2/22/83	
Per	IB	5,147,877	9/15/92	Got	ılet, Ma	rk			9/12/91	
	FOREIGN PATENT DOCUMENTS									
JUN 0 4 20	1 2	Document Number	Date		Country		Class	Sub- Class	Trans- lation	
THADEWAS	Ус	EP-196841	10/8/86	ÉΡ	EPO		1		Yes	
TABLE	ID	DE3508251	9/11/86		Germany				No	
	IE	E/P88350	9/14/83	EP	<u> </u>				Yes	
Lh_	IF	EP73143	3/2/83	Εľ	ΕΡΌ				Yes	
7	IG	Soai, Kens Functional allylation esters: co coordinat: 3290-5. (1	so and Islized term of chiral of chiral of ion of Leville 1888	hizaki, tiary a al α-ke sterec	Miyuki, lcohols to amide	, "Asyr by dia es deri	mmetri astere ived f	c Synt oselec rom (S satura	hesis of tive)-proline ted	
	IH	Soai, Kens of α-hydro chiral α-l presence o	so et al. oxy acids	, "Asym by the es with l salt,	metric s diaster complex Chem.	synthes reosele k metal Lett.,	sis of ective l hydr , 1986	both reduc ides i	eaniomers tion of n the 1897-900.	
	II	Soai, Kens reduction esters wit active α-l	so and Ha of chira th sodium hydroxy a	segawa, l α-ket borohy cids,"	Hitoshi oamides dride. I J. Chem	i, "Dia derive Prepara . Soc.,	astere ed fro ation , 1985	oselec m (S)- of opt , 1(4)	tive proline ically , 769-72.	
\sim	IJ	Soai, Kens asymmetric allyltrime alcohols,	etnvisila	ne. Pr	eparatio	on or r	protec	tea no	ive moallylic	
	IK									
N	1 1 1 2 7 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2									
Examiner		1 Cook		Date	Consider	ed:///	01			
Examiner Corl Date Considered:										

Attorney Docket Serial Number 22789XA-T 09/781,427 FORM PTO-1449 Applicant STEINER et al. INFORMATION DISCLOSURE CITATION Filing Date Feb. 13, 2001 Group Art Unit 1614 U.S. PATENT DOCUMENTS Examiner Document Sub-Filing Date Name Class Initial Number Class Date 5/20/97 JA 5,631,017 Sharpe, et al. 3/26/93 12/30/97 ιTR 5,703,088 Sharpe, et al. 6/4/92 FOREIGN PATENT DOCUMENTS JUN 0 4 2001 Document Sub-Trans-Country Date Class Number Class lation Ēρ BP--50800 5/5/82 JC EBÓ. Yes PADEW FP--48159 3/24/82 E/FIO JD Yes EF 6/25/80 EYO JΕ EP--12401 Yes OTHER (Including Author, Title, Date, Pertinent Pages, Colombo, L. et al., "Enantioselective synthesis of secondary alcohols in the presence of chiral ligands," JF Tetrahedron, 1982, 38(17), 2725-7. Soai, Kenso et al., "Unusual effect of a mixed solvent on the asymmetric reduction of chiral α-keto amides with sodium borohydride," J. Chem. Soc., 1982, 21, 1282-3. JG Steglich, Wolfgang et al., "Activated carboxylic acid derivatives. II. A simple synthesis of 2-oxycarboxylic acid amides, N-(2-oxoacyl)amino acid esters and 2-oxocarboxylic JH acid hydrazides, "Synthesis, 1978, 8, 622-4. (German) Cushman, D.W. et al., "Design of potent competitive inhibitors of angiotensin-converting enzyme. Carboxyalkanoyl and mercaptoalkanoyl amino acids," JΙ Biochemistry, 1977, 16(25), 5484-91. Steglich, Wolfgang and Hinze, Sabine, "A rational synthesis of N-trifluroacetylamino acids," Synthesis, 1976, 8, 399-JJ 401. (German) JΚ Marshall, J.A. et al., "Convenient synthesis of dioxopiperazines via aminolysis of .alpha.-(pyruvylamino) esters, Synth. Commun., 1975, 5(3), 237-44. 1/ JLHaeusler, Johannes and Schmidt, Ulrich, "Amino acids and peptides. IX. Pyruvoyl amino acids," Chem. Ber., 1974, 107(1), 145-51. (German) JM Hearn, Walter R., and Worthington, Robert E., "L-Proline-Noxalic anhydride," J. Org. Chem., 1967, 32(12), 4072-4. JN Date Considered 9/11/01 Examiner wh EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP § 609. Draw line through citation if not in conformance and not considered.

FORM PTO-1449 INFORMATION DISCLOSURE CITATION

Attorney Docket Serial Number 22789XA-T 09/781,427

Applicant STEINER et al.

Filing Date Group Art Unit Feb. 13, 2001 1614

			U.S. PAT	ENT DOCUMENTS			· · · · · · · · · · · · · · · · · · ·	
Examiner Initial		Document Number	Date	Name	Class	Sub- Class	Filing Date	
1	KA	5,424,454	6/13/95	Burbaum, B.W. et al.			5/26/94	
W. K.	\кв	5,319,098	6/7/94	Burbaum, B.W et al.			5/18/93	
JUN 0 4 2001	131		FOREIGN PA	ATENT DOCUMENTS				
. હે	<i>y</i>	Document Number	Date	Country	Class	Sub- Class	Trans- lation	
MADEMARKO	KC	WQ9200278	1/9/92	WU PCT		,	Yes	
1	KD	WQ9606097	2/29/96	·WO PCT,			Yes	
	KE	WO9512572	5/11/95	WO POT/			Yes	
	KF	WQ9407858	4/14/94	WU POT		\	Yes	
	KG	WФ9325546	12/23/93	WO PC/T		Х	Yes	
	KH	W09313066	7/8/93	WO P¢T		/ \	Yes	
	KI	J₽05178824	7/20/93	JP Japan	/		No	
	KJ	EP-572365	12/1/93	₽Ø EPd			Yes	
-	KK	WO9219593	11/12/92	W) PCT			Yes	
	KL	GB2247456	3/4/92	6 United Kingdom	/		Yes	
W	KM	WØ9218478	10/29/92	W) Pat	Z	,	Yes	
	OTHE	R (Including A	uthor, Tit	le, Date, Pertinent	Pages,	etc.)		
N	. KN	Chakarabo of cyclic	rty, Tush	ar K., "Studies towa based analogs of mac ," Pure Appl. Chem.,	rds the	e deve	lopment	
	ко							
	KP							
if	KQ	Tugwell, Rheumatoi	Peter, "C d Arthrit	lyclosporin in the T is," J. of Autoimmun	reatmenty, 19	nt of 992, 5	, 231-40.	
Fry, Lionel, "Psoriasis: Immunopathology and Long-term treatment with Cyclosporin," J. of Autoimmunity, 1992, 5, 277-83.								
h	KS	Feutren, Autoimmun	Gilles, " e Disease	The Optimal use of C s," J. of Autoimmuni	Ly, IJ.	orin A 92, 5,	in 183-95.	
Examiner	A	N Cerch		Date Considered:	1			

EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP § 609.

Draw line through citation if not in conformance and not considered.

FORM PTO-1449

INFORMATION DISCLOSURE CITATION

Attorney Docket 22789XA-T

Serial Number 09/781,427

Applicant

STEINER et al.

Filing Date Feb. 13, 2001 Group Art Unit 1614

Feb. 13, 2001 1614									
			U.S. PAT	ENT DO	CUMENTS				
Examiner Initial		Document Number	Date		Name	Class	Sub- Class	Filing Date	
010	ĽΑ	5,614,547	3/25/97	Hami	lton et. al			6/7/95	
6.7	LB								
UN 0 4 2007 S		E	OREIGN PA	ATENT D	OCUMENTS				
ette /		Document Number	Date		Country	Class	Sub- Class	Trans- lation	
HOEMARK	LC	EP-652229	5/10/95	ļ	EPØ			Yes	
<u> </u>	LD	Z2•9207782	4/28/93	ZA Sot	th Afric a			Yes	
	OTHER	R (Including Au	thor, Tit	le, Da	te, Pertinent	Pages,	etc.)		
^	LE	HIV and F of Pyrrol	IV Protea idine-Con hylamine	se: Inh taining Core St	"Selectivity nibitory and M g α-Keto Amide tructures, J.	echani and	stic S	tudies	
	LF	Nicolaou, CheEur	K.C. et 1, 199	al., "1 5 , 1(5)	Total synthesi , 318-33:	s of r	apamyc	<u>in</u> ,"	
W.	LG	new core	structure	for th	Ketoamide Phe ne inhibition 2(10), 1085-	ot HIV	soster prote	e as a ase,"	
	LH	Hauske, James R. et al., "Investigation of the effects of							
	LI	butylamin carboxype angiotens	opropoxy) ntyl)-DL- in-conver cking pro	<pre>-indole alanyl] ting er</pre>	"1-[4-(2-Hyd e-3-yl (5-acet -L-proline di azyme inhibito s," KhimFarm	amido- hydroc r with	1-(S)- hlorid β-	e, a new	
	LJ	Ranganath Modificat Am. Chem.	ion by No	vel Cα-	al., "Protein : C Side-Chain : 6545-57.	Backbo: Scissi	ne on," 1	994, J.	
	LK	isolated	microsome	s and i	"Inhibition o no acid deriv n embryonic c , 525-30.	f prolatives	yl 4- in vi tissu	tro, in es,"	
	LL	Increases	the Rate	of Axo	The Immunosup nal Regenerat 1995, 15(11):	ion in	Rat S	06 ciatic	
	Karle, Isabella L. et al., "Coformation of the oxalamide group in retro-bispeptides. Three crystal structures," Int. J. Pept. Protein Res., 1994, 43(2), 160-5.								
N	LN	Kaczmar, (German).	et al., M	akromol	. Chem., 1976	, 177,	1981-	9	
Examiner		A Cook	/		Date Considere				
		,							

EXAMINER: I

Initial if reference considered, whether or not citation is in conformance with MPEP § 609. Draw line through citation if not in conformance and not considered.

OIP				· .			Page	13 of 2
JUN 0 4 2001	yC131	FORM PTO-144	^	-"-	Attorney Dock 22789XA-T		rial N 09/78	umber 31,427
e .	&/			,	Applicant			· · · · · · · · · · · · · · · · · · ·
PADEMARBON	ORMA	TION DISCLOSUR	E CITATIO	<u>N</u>	Filing Date	INER e		
					Feb. 13, 20		-	t Unit 514
			U.S. PAT	ENT DO	CUMENTS			
Examiner Initial		Document Number	Date		Name	Class	Sub- Class	Filing Date
	MA							
	MB	<u> </u>						
		I	OREIGN PA	ATENT D	OCUMENTS			
		Document Number	Date		Country	Class	Sub- Class	Trans- lation
Л	MC	₩ 0 9405639	3/17/94	Μó	PCT	\rightarrow	<u> </u>	Yes
<u> </u>	MD	WO9615101	5/23/96	NO	PCT		<u> </u>	Yes
	OTHE	R (Including Au	thor, Tit	le, Da	te, Pertinent	Pages,	etc.)
n	ME	Steiner, immunophi Lett., 19	lin FKBP	coloca	., "High brain alized with ca	densi lcineu	ties c rin,"	f the Nature
	MF	the trica	rbonyl su	bunit :	nkard, Mark, " in the immunos Lett., 1993, 3	uppres	sant	
	MG	Furber, M immunosup 1993, 34(pressive	activit	ies relating t tiy of FK506,"	o the Tetra	hedron	Lett.,
	МН	Ranganath for prote	an, Darsh in design	an et a	al., "Oxalopep Chem. Soc., 19	tides 93, (1	as cor), 92-	e motifs 4.
	MI	phosphory	lation of lutamate	nitrio neuroto	Immunosuppress c oxide syntha oxicity," Proc	se and	prote	cts
~	MJ	Cunliffe, hydroxyla oxaloglyc (14), 265	ine and i	et al. Inhibit ts der:	"Novel inhib tion by the su lvatives," J.	oitors bstrat Med. C	of pro e anal hem.,	olyl 4- og N- 1992, 35
	MK	Llauxiliari	es in Bar	bier-ti	pacid esters pe reactions Chem., 1991,	in aqu	eous	2.
N	ML	on the bi	ological	activit	ct of the natu ty of N-carbox 5(7), 44-6. (R	yalkyl	dipep	radical tides,"
	MM	Blaschke	e t al., C	hemica	Abstracts, 1	974, 0	5:7840	5k.
Examiner		n Corh			Date Consider	ed (01		

Initial if reference considered, whether or not citation is in conformance with MPEP § 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to Applicant. EXAMINER:

ORMATION DISCLOSURE CITATION

FORM PTO-1449

Attorney Docket 22789XA-T

Serial Number 09/781,427

Applicant

STEINER et al.

Filing Date Feb. 13, 2001

Group Art Unit 1614

				Feb. 13, 2	2001	<u></u>	74			
U.S. PATENT DOCUMENTS										
Examiner Initial		Document Number	Date	Name	Class	Sub- Class	Filing Date			
	NA	5,414,083	5/9/95	Hackl et al.	1	/	1/24/94			
	NB	4,310,461	1/12/82	Krapcho et al.			1/23/80			
	NC	4,578,474	3/25/86	Krapcho et al.			11/19/84			
i\	ND	4,531,964	7/30/85	Shimano et al.)	8/29/83			
		F	FOREIGN PA	ATENT DOCUMENTS						
	Document Number Date Country Class Sub- Trans- Class lation									
V	NE	₩ 0 9535367	12/28/95	¡No PCT			Yes			
(A /	NF	₩ 09 413629	6/23/94	WU PCT		/	Yes			
	OTHE	R (Including Au	thor, Tit	le, Date, Pertiner	ıt Pages,	etc.)				
W_1	NG	Caufield, Immunomod Johns (Ed	Craig E. ulators," .), Acade	and Musser, John <u>Annual Reports in</u> mic Press, Chapter	H., "Mac <u>Medicin</u> 21, 195	rocycl al Che -204,	ic mistry, 1989.			
	NH	Effenberg benzenesu	er F. et lfenvl ch	al., "Diastereosel loride to 1-acrylo , 1989, 110:154846	ective a vlprolin	dditio	n of			
N	NI	Nakatsuka Reagent, 112 (14),	, M et al $(C_8, C_9 - 1)$ 5583-90.	. " Total Synthesi ³ C ₂)-FK-506," J. Am ·	s of FK5 . Chem.	06 and Soc.,	an FKBP 1990,			
	ŊĴ	Shu, A. e rapamycin 38(3), 27	t al., "S analogs,	ynthesis of I-125. " J. Labelled Comp	labeled d. Radio	photoa pharm.	ffinity , 1996,			
7	Tatlock, J. et al., "High affinity FKBP-12 ligands from									
	NL	Teague, S macrocycl 6.	. et al., es," Bioo	"Synthesis of FK5 rg. Med. Chem. Let	06-cylco t., 1995	sporin , 5(20	hybrid), 2341-			
W	NM	Stocks, M the intra 1995, 36(molecular	"Macrocyclic ring Heck reaction," T -8.	closure etrahedr	s empl on Let	oying t.,			

Examiner

Date Considered

Initial if reference considered, whether or not citation is in conformance with MPEP § 609. Draw line through citation if not in conformance and not considered.

Include copy of this form with next communication to Applicant.

JUN 0 6 2001

TECH CENTER 1600/2900

Examiner Initial	K 10131	FORM PTO-144: TION DISCLOSURE Document Number	E CITATION	Attorney Do 22789XA Applicant S Filing Date Feb. 13, 2 ENT DOCUMENTS Name	TEINE	R e	t al. oup Art 16	1,427		
4	OA	4,390,695	1/28/83	Krapcho et al.				6/1/81		
1	ОВ	4,593,102	6/3/86	Shanklin, Jr.		<u>/</u>	>	7/1/95		
	OC									
	OD									
]	FOREIGN PA	TENT DOCUMENTS						
		Document Number	Date	Country	Cl	ass	Sub- Class	Trans- lation		
\wedge	OE	₩ 0 9204370	3/19/92	WO PCT		\searrow		Yes		
\sim	OF	₩09535308	12/28/95	WU PCT		_		Yes		
M	Wang, C.P. et al., "High performance liquid chromatographic isolation and spectroscopic characterization of three major metabolites from the plasma of rats receiving rapamycin (sirolimus) orally," J. Liq. Chromatogr., 1995, 18(13), 2559-68. Armistead, D.M. et al., "Design, synthesis and structure of non-macrocyclic inhibitors of FKBP12, the major binding protin for the immunosuppressant FK506," Acta Crystallogr. 1995, D51(4), 522-8. Luengo, J. et al., "Structure-activity studies of rapamycir analogs: evidence that the C-7 methodoxy group is part of the effector domain and positioned at the FKBP:12-FRAP interface," Chem. Biol., 1995, 2(7), 471-81. Furber, Mark, "FKBP-12-ligand-calceineurin interactions:									
~	OK Chakraborty, TK et al., "Design and Synthesis of a rapamycin-based high affinity binding FKBP12 ligand," Chem. Biol., 1995, 2(3), 157-61. Wang, C.P. et al., "A high performance liquid chromatographic method for the determination of rapamycin {sirolimus} in rat serum, plasma, and blood and in monkey serum," J. Liq. Chromatogr., 1995, 18(9), 1801-8.									
	Smith, A.B. et al., "Total synthesis of rapamycin and demethoxyrapamycin," J. Am. Chem. soc., 1995, 117(19), 5407-8. Date Considered Aminer: Initial if reference considered, whether or not citation is in conformance with MPEP § 609. Draw line through citation if not in conformance and not considered.									

JUN 0 4 2001

FORM PTO-1449

MATION DISCLOSURE CITATION

Attorney Docket 22789XA-T Serial Number 09/781,427

Applicant

STEINER et al.

Filing Date Feb. 13, 2001 Group Art Unit 1614

			U.S. PAT	ENT DOCUMENTS							
Examiner Initial		Document Number	Date	Name	Class	Sub- Class	Filing Date				
	PA										
	PB										
	FOREIGN PATENT DOCUMENTS										
		Document Number	Date	Country	Class	Sub- Class	Trans- lation				
\mathcal{A}	PC	₩09221313	12/10/92	WO PCT	\longrightarrow	<u> </u>	Yes				
4	PD	WO9524385	9/14/95	WO PCT			Yes				
	OTHE	R (Including Au	thor, Tit	le, Date, Perti	nent Pages,	etc.)					
2	PE	the major	equilibr	, "Synthesis an ium products of 1995, 26(13),	ascomycin	cleav and Fk	age of 506,"				
	PF	Nelson, F rapamycin	Nelson, F. et al., "A novel ring contraction of rapamycin," Tetrahedron Lett., 1994, 35(41), 7557-60.								
	PG	Dawson, T cyclophil relations 569-80.	Dawson, T.M. et al., "The immunophilins, FK506 binding and cyclophilin, are discretely localized in the brain: relationship to calcineurin," Neuroscience, 1994, 62(2), 569-80.								
	PH	receptor	Cameron, Andrew et al., "Immunophilin FK506 binding protein associated with inositol 1,4,5-triphosphate receptor modulates calcium flux," Proc. Natl. Acad. Sci. USA, 1995, 92, 1784-1788.								
	ΡI	Stocks, M pyranosid FK-506,"	Stocks, M. et al., "The contribution to the binding of the pyranoside sustituents in the excised binding domain of FK-506," Bioorg. Med. Chem. Lett., 1994, 4(12), 1457-60.								
	PJ	Steiner, J.P. et al., "Nonimmunosuppressive Ligands for Neuroimmunophilins Promote Nerve Extension <i>In Vitro</i> and <i>In Vivo</i> ," Society for Neuroscience Abstracts, 1996, 22, 297.13.									
	PK	lthe Expre	Lyons, W. Ernest et al., "Neronal Regeneration Enhances the Expression of the Immunophilin FKBP-12," The Journal of Neuroscience, 1995, 15, 2985-94.								
	PL	Skotnicki derivativ	y Jerauld es," Tetr	et al., "Ring ahedron Lett.,	expanded ra 1994, 35(2)	pamyci , 201-	n 2.				
N	PM	Skotnicki esters an	, Jerauld d amides,	et al., "Synth " Tetrah. Lett.	esis of sec , 1994, 35(orapam 2), 19	ycin 7-200.				
Data Congidered(

Examiner

R Corh

Date Considered

9/10/ECEIVED

EXAMINER: Inif reference considered, whether or not citation is in conformance with MPEP \$ 609.

Draw line through citation if not in conformance and not considered.

Include copy of this form with next communication to Applicant itial



Page 17 of 21 Attorney Docket Serial Number 22789XA-T 09/781,427 FORM PTO-1449 Applicant STEINER et al. MATION DISCLOSURE CITATION Filing Date Group Art Unit Feb. 13, 2001 1614 U.S. PATENT DOCUMENTS Sub-Filing Examiner Document Date Name Class Initial Number Class Date OA OB FOREIGN PATENT DOCUMENTS Document Sub-Trans-Date Country Class Number Class lation 6M QC W09307269 4/15/93 PCT Yes 10/1/92 PCT wJ OD WO9216501 Yes (Including Author, Title, Date, Pertinent Pages, etc.) OTHER Rao, A.V. Rama and Desibhatla, Vidyanand, "Studies directed towards the syntesis of rapamycin: OE stereoselective synthesis of C-1 to C-15 segment," Tetrahedron Lett., 1993, 34(44), 7111-14. Andrus, Merrit B., "Structure-based design of an acyclic ligand that bridges FKBP12 and calcineurin, " J. Am. Chem. OF Soc., 1993, 115(2), 10420-1. Luengo, Juan I. et al., "Efficient removal of pipecolinate from rapamycin and FK506 by reaction with OG tetrabutylammonium cyanide, "Tetrahedron Lett., 1993, 34(29), 4599-602. Steffan, Robert J. et al., "Base catalyzed degradations of NI OH rapamycin, Tetrahedron Lett., 1993, 34(23), 3699-702. Nicolaou, K.C. et al., "Total Synthesis of rapamycin," J. OI Am. Chem. Sec., 1993, 115(10), 4419-20. Hayward, C.M. et al., "Total Synthesis of rapamycin via a U novel titanium-mediated aldol macrocyclization reaction," OJ J. Am. Chem. Soc., 1993, 115(20), 9345-6. Yohannes, Daniel et al., "Degradation of rapamycin: synthesis of a rapamycin-derived fragment containing the OK tricarbonyl and triene sectors, "Tetrahedron Lett., 1993, 34(13), 2075-8. Luengo, J. et al., "Studies on the chemistry of rapamycin: novel transformation under Lewis-acid catalysis," QL N/

Tetrahedron Lett., 1993, 34(6), 991-4.

Examiner

Date Considered

Initial if reference considered, whether or not citation is in conformance with MPRP § 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to Applicant.

010							Page	18 of 21		
JUN 0 4 2001	40	FORM PTO-144	•		Attorney Dock 22789XA-T	et Se	rial N 09/78	umber 31,427		
Yllae	Ju/	TION DISCLOSUR		NT.	Applicant STEINER et al.					
PADEMAR	~	TION DIBELLOSON	<u> </u>	<u> </u>	Filing Date Feb. 13, 200			t Unit 514		
			U.S. PAT	ENT DO	CUMENTS					
Examiner Initial		Document Number	Date		Name		Sub- Class	Filing Date		
	RA RB									
	KB]	F	OREIGN PA	ר ייאייי	OCUMENTS	<u>i</u>	<u> </u>			
		Document	Date		Country	Class	Sub-	Trans-		
	RC	Number - \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	10/5/95	พง	PET		Class	lation Yes		
	OTHER			-	te, Pertinent	Pages	etc \			
		Yohannes,	Daniel e	t al.,	"Degradation	of rap	amycin	:		
7	RD	retrieval 1992, 33(of major	intact	subunits," T	etrahē	dron L	ett.,		
	RE	Goulet, M the trica Tetrahedr	Goulet, Mark T. and Boger, Joshua, "Degradative studies on the tricarbonyl containing macrolide rapamycin," Tetrahedron Lett., 1991, 32(45), 6454.							
	RF	from rapa	Goulet, Mark T. et al., "Construction of the FK-506 analog from rapamycin-derived materials," Tetrahedron Lett., 1991, 32(36), 4627-30.							
	RG	synthesis	Rao, A.V. Rama et al., "Studies directed towards the synthesis of immunosuppressive agent FK-506: synthesis of the entire bottom half," Tetrahedron Lett., 1991, 32(9),							
	RH	carbon-ca	rbon bond	cleava	On the remark age reactions g. Chem., 1991	in teh	C(8)-	C(10)		
	RI	Linde, Ro 1,2,3-tri 2534-8.	bert G. e carbonyl	t al., systems	"Straightforw s," J. Org. Ch	ard sy em., 1	nthesi 991, 5	s of 6(7),		
	RJ	Hayward, reaction segment o	C.M. et a to the re f rapamyc	l., "Ar giospec in," 39	application cific synthesi 989-92.	of the s of t	Suare he C ₂₈ -	Z C ₄₂		
	RK	Hovarth, R., et al., "An application of the Evans-Prasad 1,3-Syn diol synthesis to a stereospecific synthesis of the C_{10} - C_{27} segment of rapamycin," Tetrahedron Lett., 1993, 34(25), 3993-3996.								
	RL	Whitesell, J.K. et al., "Asymmetric Induction. Reduction, Nucleophilic Addition to, Ene Reactions of Chiral α-Ketoesters," J. Chem. Soc., Chem Commun., 1983, 802.								
	RM	Ando, Takao et al., "Formation of Crossed Phenzine from								
Examiner		a Corh			Date Considere	/ A	JN 0 6	2001		
EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP § 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to Applicant. TECH CENTER 1600/2000										



JUN 0 4 2001 STATE AND EMARKS

FORM PTO-1449

TION DISCLOSURE CITATION

Attorney Docket 22789XA-T

Serial Number 09/781,427

Applicant

STEINER et al.

Filing Date Feb. 13, 2001 Group Art Unit 1614

NOEW			and the second		Feb. 13,		16	14
			U.S. PAT	ENT DOC	UMENTS			
Examiner Initial		Document Number	Date		Name	Class	Sub- Class	Filing Date
		I	OREIGN PA	TENT D	OCUMENTS		•	
		Document Number	Date		Country	Class	Sub- Class	Trans- lation
N	SA	WO9219745	11/12/92	W o	Р¢т			Yes
	SB	WO9323548	11/25/93	Wo	PCT		+/-	Yes
	sc	WD9636630	11/21/96	WO	РСТ	\rightarrow		Yes
	SD	WD9633187	10/24/96	ΝJ	РСТ		<u>//</u>	Yes
	SE	WD9633184	10/24/96	WO	РСТ		V	Yes
	SF	WD9603318	10/24/96	W	РСТ	/	1	Yes
	SG	WD9820891	5/22/98	WØ	РСТ		\	Yes
	SH	WD9820892	5/22/98	Wo	PCT	/_	1	Yes
	SI	WD9820893	5/22/98	W	PCT_	/	$\bot ackslash$	Yes
\sim	SJ	WO9824805	6/11/98	W l	РФТ			Yes
	OTHE	R (Including A	uthor, Tit	le, Da	te, Pertin	ent Pages	, etc.)	
. W	SK	Kino, Tor isolatede 40(9), 12	ed from A	"FK-50 strepto	omyces," J	l immunosu . of Antik	ippress piotics	nt , 1987,
	SL	Waldmann, auxilary 1990, Syn	Herbert, im Barbie ilett, 10,	r type 627-8.		in aqueoi	ıs solu	tion,"
N	SM	Neurodege Sci. USA,	nerative 94:2019-	Animal 2024.	., "Neuro iral and Fi Models,"	1997, Prod	c. Nati	. Acea.
N	SN	Steiner, Nonimmund FK506, Ra 428.	Joseph P. suppressi pamycin a	, et alve Analnd Cycl	., "Neuro Logues of Losporin A	trophic Ad Immunosupp ," Nat. Me	ctions pressived. 3(4	of e Drugs):421-
	so							
	SP						문 오	교
Examiner		R Ca	rh		Date Cons	idered 9///0/	CENTER 1	Z C
Draw	line throu	erence considered, whether or	rmance and not cons	idered.	with MPEP § 609.		H (N Z
Inclu	ae copy of	this form with next commun	reaction to Applican				8	<u> </u>

70 11 & 3	63		•	Attorney Docket Serial Numbe 22789XA-T 09/781,42						
JUN 0 4 2001) (3)	FORM PTO-144		_	Applicant STEINER et al.					
I NIP	ØRMA'	rion disclosur	E CITATION		Filing Date Group Art Unit Feb. 13, 2001 1614					
U.S. PATENT DOCUMENTS										
Examiner Initial					Name C		Sub- Class	Filing Date		
	_									
				<u> </u>			-			
					 	<u></u>				
					O CURATION C	<u> </u>				
		Document	FOREIGN PA		Sub- Tr			Trans-		
		Number	Date		Country	Clas	SS Class	lation		
		- M								
		<u> </u>				<u> </u>				
	OTHE	R (Including A								
	TA	Immunoph:	ilin Ligan	ds, Am	Hair Growth Mc . J. Path. 150	:143	3-41 (19	97).		
										
Examiner	Examiner R $Considered$ $g/(/a)$									
Draw 1	design in the supplier with MDPD 5 600									



FORM PTO-1449

ORMATION DISCLOSURE CITATION

Attorney Docket 22789XA-T

Serial Number 09/781,427

Applicant

STEINER et_al.

Filing Date Feb. 13, 2001

Group Art Unit 1614

				reb. 13, 20	<u> </u>)14
		<u> </u>	U.S. PAT	ENT DOCUMENTS			
Examiner Initial		Document Number	Date	Name	Class	Sub- Class	Filing Date
W	UA	5,714,510	2-3-98	Proctor			6-5-95
, and the second	UB	5,620,971	4-15-97	Armistead et al.			3-25-94
	UC	5,472,687	12-5-95	Proctor		X	2-7-94
	UD	5,385,908	1-31-95	Nelson et al.			11-22-93
	UE	4,438,031	3-20-84	Winkley et al.		T	2-24-82
		1	FOREIGN PA	ATENT DOCUMENTS		-	
		Document Number	Date	Country	Class	Sub- Class	Trans- lation
	UF	DF 2505114	8-19-76	Ò€ Germany)	NO
	UG	EP 0823419	8-7-97	ЕРΦ			YES
	UH	EP 0519819	6-17-92	ЕРФ			NO
	UI	EP 0494005	12-20-91	EPO			NO
	UJ	EP 0471135	8-14-90	EPO			YES
	UK	EP 0443983	12-2-91	ЕРФ			NO
	UL	EP 0420707	8-24-90	ЕРФ			NO
	UM	WO 9822432	5-28-98	WO			NO
	UN	WD 9813343	4-2-98	WU PCT			YES
	UO	WD 9731898	9-4-97	wo ect			YES
	UP	WD 9611943	10-6-95	wo por			YES
	UQ	WO 9534303	12-21-95	WU POT			YES
	UR	WO 9512398	5-11-95	WO ECT			YES
	US	WD 9502684	1-26-95	WO PCT			YES
	UT	WO 9403476	2-17-94	WV PCT			YES
	UU	WO 9318736	9-30-93	WO PAT			NO
	UV	WO 9314762	8-5-93	WO POT			YES
	UW	WO 9314072	7-22-93	NO PCT			YES
	UX	WO 8906234	7-13-89	WO PCT			YES
\mathcal{N}	UY	WO 8800040	1-14-88	WO PGI		\	YES

Examiner

Date Considere RECEIVED

Initial if reference considered, whether or not citation is in conformance with MPEP \$ 609. Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to Applicant.